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ABSTRACT

BACKGROUND AND PURPOSE: *Isocitrate dehydrogenase (IDH)* has been shown to have both diagnostic and prognostic implications in gliomas. The purpose of this study was to examine whether DWI and DSC-PWI combined with conventional MR imaging could noninvasively predict *IDH* mutational status in World Health Organization grade II and III astrocytomas.

MATERIALS AND METHODS: We retrospectively reviewed DWI, DSC-PWI, and conventional MR imaging in 42 patients with World Health Organization grade II and III astrocytomas. Minimum ADC, relative ADC, and relative maximum CBV values were compared between *IDH*-mutant and wild-type tumors by using the Mann-Whitney *U* test. Receiver operating characteristic curve and logistic regression were used to assess their diagnostic performances.

RESULTS: Minimum ADC and relative ADC were significantly higher in *IDH*-mutated grade II and III astrocytomas than in *IDH* wild-type tumors ($P < .05$). Minimum ADC with the cutoff value of $\geq 1.01 \times 10^{-3} \text{ mm}^2/\text{s}$ could differentiate the mutational status with a sensitivity, specificity, positive predictive value, and negative predictive value of 76.9%, 82.6%, 91.2%, and 60.5%, respectively. The threshold value of < 2.35 for relative maximum CBV in the prediction of *IDH* mutation provided a sensitivity, specificity, positive predictive value, and negative predictive value of 100.0%, 60.9%, 85.6%, and 100.0%, respectively. A combination of DWI, DSC-PWI, and conventional MR imaging for the identification of *IDH* mutations resulted in a sensitivity, specificity, positive predictive value, and negative predictive value of 92.3%, 91.3%, 96.1%, and 83.6%.

CONCLUSIONS: A combination of conventional MR imaging, DWI, and DSC-PWI techniques produces a high sensitivity, specificity, positive predictive value, and negative predictive value for predicting *IDH* mutations in grade II and III astrocytomas. The strategy of using advanced, semiquantitative MR imaging techniques may provide an important, noninvasive, surrogate marker that should be studied further in larger, prospective trials.

ABBREVIATIONS: ADC_{min} = minimum ADC; cMRI = conventional MR imaging; *IDH* = *isocitrate dehydrogenase*; rADC = relative ADC; rCBV = relative CBV; rCBV_{max} = relative maximum CBV; WHO = World Health Organization

Infiltrating astrocytomas are the most common primary central nervous system tumors, ranging variably from grade II to IV according to the 2007 World Health Organization classification

system.^{1,2} Glioma grading is based on histopathologic analysis of tumor differentiation, mitotic activity, cellularity, nuclear atypia, and the extent of microvascular proliferation and may result in a great deal of interobserver variability.¹⁻³ Therefore, quantitative molecular analyses have the potential to reduce subjectivity and improve diagnosis, prognostication, risk stratification, and management plans. Notably, in the 2016 World Health Organization (WHO) classification, grade II and III astrocytomas are molecularly divided into *isocitrate dehydrogenase (IDH)* mutant, *IDH* wild type, and not otherwise specified categories, emphasizing the value of *IDH* mutation status in astrocytomas.⁴

IDH gene mutations, originally discovered in high-grade gliomas in 2008, exist in 60%–90% of grade II and III astrocytomas.^{5,6} The *IDH* gene (including *IDH1* and *IDH2* genes) plays prominent roles in the metabolism, pathogenesis, and progression of astrocytomas.⁷⁻⁹ In addition, stratification of grade II and III gliomas

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into subsets defined by *IDH* mutation would help identify subgroups with distinct prognostic characteristics, therapeutic response, and clinical management.^{10–16} In a study with a cohort of 475 patients, comparison of overall survival between those with WHO grade II and III *IDH*-mutated astrocytomas showed no remarkable difference, whereas the patients with *IDH*-mutated tumors survived much longer than those with *IDH* wild-type tumors.¹⁰ Patients with grade II astrocytomas without *IDH* mutation were shown to have a poorer prognosis with a 5-year progression-free survival and overall survival rate of 14% and 51%, respectively, compared with 42% and 93% for those with *IDH*-mutant tumors ($P < .001$).¹⁷ Moreover, patients with gliomas whose lesions had *IDH* mutations were more sensitive to chemoradiation therapy and survived longer than those with wild-type *IDH*.^{10,15}

Currently, immunohistochemical staining and DNA sequencing are the most common methods for determining the *IDH* mutational status in gliomas. *IDH* gene mutations may reflect alterations in metabolism, cellularity, and angiogenesis, which may manifest characteristic features on DWI and DSC-PWI.^{8,18} DWI can noninvasively provide direct insight into the microscopic physical properties of tissues through observing the Brownian movement of water and reflecting cellularity within the lesions by ADC values.^{19–21} In vivo measurement of relative CBV (rCBV) has been demonstrated to correlate with tumor vascularity.^{22–24} ADC values derived from DWI and DSC-PWI have been used to detect *IDH* gene status in gliomas in recent research.^{24–26} Meanwhile, conventional MR imaging (cMRI) was also used to assess other characteristics of gliomas (eg, location, distinctness of borders, enhancement, and edema).^{17,27,28}

To our knowledge, there is no study in the literature combining cMRI with diffusion and perfusion techniques to distinguish *IDH* genotypes. The purpose of this study was to explore whether a novel approach, in which DWI and DSC-PWI were combined with cMRI, was able to noninvasively predict *IDH* mutational status in WHO grade II and III astrocytomas.

MATERIALS AND METHODS

Patients

The ethics committee of our hospital approved this retrospective study, and the requirement for patient informed consent was waived due to the nature of the retrospective study. Ninety-six patients who underwent surgical resection or stereotactic biopsy at our institution from July 2014 through June 2016 were selected. The inclusion criteria were as follows: 1) definite histopathologic diagnosis of grade II and III astrocytomas based on the WHO 2016 classification criteria, 2) cMRI, DWI, and DSC-PWI performed before treatment, and 3) all data available in 3T MR imaging. As a result, 42 patients (26 males and 16 females; mean age, 41.83 ± 15.98 years; age range, 8–72 years) were included in the study.

MR Imaging Techniques

Images were acquired in the routine clinical work-up on a 3T MR imaging system (Magnetom Verio Tim; Siemens, Erlangen, Germany) with an 8-channel head matrix coil. The conventional MR imaging protocols consisted of the following sequences: axial T1-

Table 1: The main clinical and cMRI features of *IDH* mutational status in grade II and III astrocytomas^a

	<i>IDH</i> Mutation (n = 17)	<i>IDH</i> Wild Type (n = 25)	P Value
Sex (male/female)	9/8	17/8	.157
Age (yr)	35.76 ± 9.13	45.96 ± 18.36	.041
Location			.006
Frontal lobe	9 (52.9%)	4 (16.0%)	
Parietal lobe	1 (5.9%)	3 (12.0%)	
Temporal lobe	6 (35.3%)	4 (16.0%)	
Occipital lobe	0	0	
Insular lobe	1 (5.9%)	3 (12.0%)	
Others	0	11 (44.0%)	
Homogeneity			.439
Homogeneous	4 (23.5%)	7 (28.0%)	
Heterogeneous	13 (76.5%)	18 (72.0%)	
Edema			.746
Presence	3 (17.6%)	7 (28.0%)	
Absence	14 (82.4%)	18 (72.0%)	
Borders			.037
Sharp	11 (64.7%)	8 (32.0%)	
Indistinct	6 (35.3%)	17 (68.0%)	
Contrast enhancement			.286
No	11 (64.7%)	12 (48.0%)	
Yes	6 (35.3%)	13 (54.0%)	
Histology			.051
Grade II astrocytomas	12 (70.6%)	12 (48.0%)	
Grade III astrocytomas	5 (29.4%)	13 (54.0%)	

^a Data are number (%) unless otherwise indicated.

weighted gradient-echo imaging (TR = 250 ms; TE = 2.48 ms), axial T2-weighted turbo spin-echo imaging (TR = 4000 ms; TE = 96 ms), axial fluid-attenuated inversion recovery imaging (TR/TE = 9000/94 ms; TI = 2500 ms), and 3 orthogonal plane contrast-enhanced gradient-echo T1-weighted imaging scans (TR/TE, 250/2.48 ms) acquired following the acquisition of DSC-PWI sequences. The section thickness (5 mm), intersection gap (1 mm), and FOV (220 × 220 mm) were uniform in all sequences.

DWI was performed in the axial plane with a spin-echo echo-planar sequence before injection of contrast material. The imaging parameters used were as follows: TR/TE = 8200/102 ms, NEX = 2.0, section thickness = 5 mm, intersection gap = 1 mm, FOV = 220 × 220 mm. The b-values were 0 and 1000 s/mm² with diffusion gradients encoded in the 3 orthogonal directions to generate 3 sets of diffusion-weighted images. Processing of the ADC map was generated automatically by the MR imaging system.

DSC-PWI was performed with a gradient-recalled T2*-weighted echo-planar imaging sequence. The imaging parameters were as follows: TR/TE = 1000–1250/54 ms, flip angle = 35°, section thickness = 5 mm, intersection gap = 1 mm, NEX = 1.0, FOV = 220 × 220 mm. During the first 3 phases, images were acquired before injecting the contrast material to establish a pre-contrast baseline. When the scan was to the fourth phase of DSC-PWI, a bolus of gadobenate dimeglumine at a dose of 0.1 mmol/kg of body weight and 5 mL/s was injected intravenously with an MR imaging-compatible power injector. After we injected a bolus of the contrast material, a 20.0-mL bolus of saline was administered at the same injection rate. The series of 20 sections, 60 phases, and 1200 images was obtained in 1 minute 36 seconds.

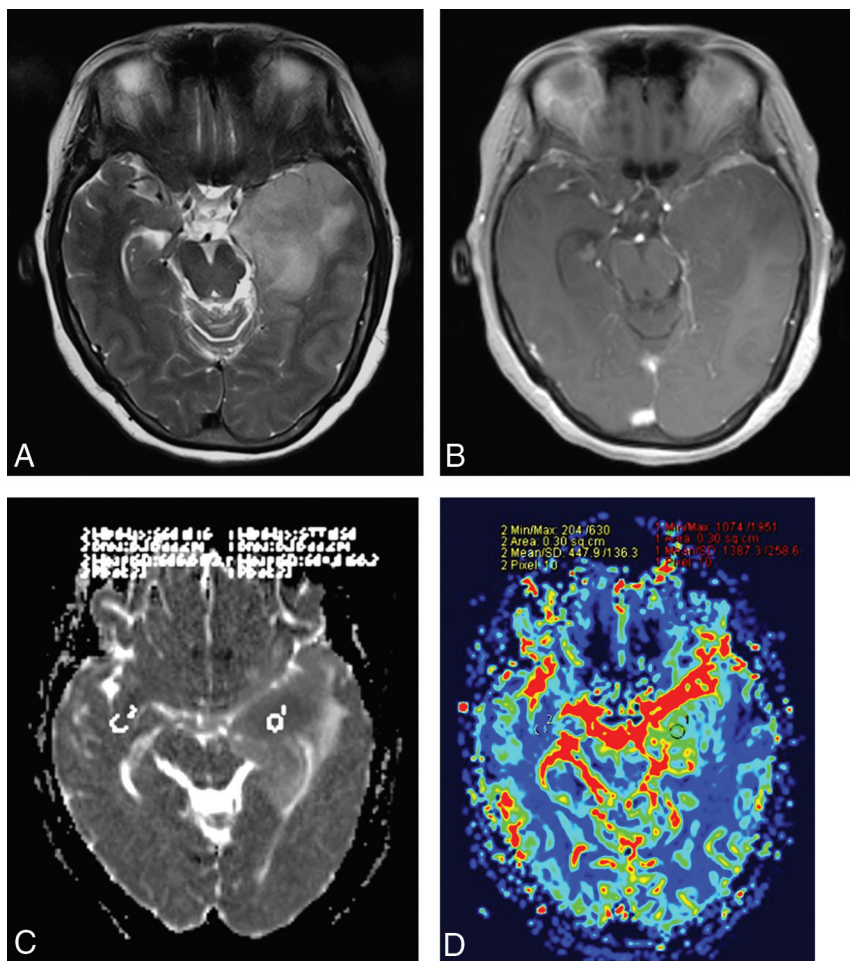


FIG 1. A 52-year-old woman with a diffuse astrocytoma without *IDH* mutation. A, Axial T2WI demonstrates heterogeneous high signal intensity with indistinct borders on the left temporal lobe. B, Contrast-enhanced axial T1-weighted image demonstrates a lesion enhancement with blurred borders. C, A corresponding ADC map shows the tumor with a decreased ADC value ($ADC_{min} = 0.684 \times 10^{-3} \text{ mm}^2/\text{s}$, $rADC = 1.08$). D, Correlative color CBV image shows elevated perfusion with the calculated $rCBV_{max}$ of 3.10.

Data Processing

Image postprocessing of perfusion data and perfusion measurements was performed on an off-line syngo B19 workstation (Siemens) with standard software. All cMRI data with respect to tumor locations, heterogeneity, borders, peritumoral edema, and contrast-enhancement pattern were assessed by 2 neuroradiologists who were blinded to tumor histology and molecular characteristics. Tumor location, considered to be the lobe within which the bulk of the tumor resided, was divided into 6 groups: frontal lobe, temporal lobe, parietal lobe, insular lobe, occipital lobe, and others (basal ganglia, thalamus, brain stem, and cerebellum). Edema was defined as a nonenhanced area on contrast-enhanced T1WI and higher signal outside the tumoral solid area on T2WI and FLAIR. Tumor borders were classified as sharp or indistinct on the basis of T2WI and FLAIR sequences (relatively decreased signal intensity on T2WI or FLAIR should be regarded as tumor area rather than edema in gliomas). A senior neuroradiologist made the final decision when 2 observers disagreed.

For evaluation of DWI data, ADC values were measured by manually selecting ROIs inside the tumor regions on the ADC maps. All continuous sections including tumors were observed.

At least 5 small round ROIs ($30\text{--}40 \text{ mm}^2$) were placed inside the tumors on the ADC maps without overlap. The bigger the tumor was, the more ROIs were selected. Finally, the ROI with the lowest ADC value was chosen to calculate minimum ADC (ADC_{min}). We made the ROI placement from the solid portion of the lesion (defined on T2WI and contrast-enhanced T1WI), avoiding hemorrhagic, cystic, necrotic, or apparent blood vessel regions that might influence the ADC values. The minimum ADC is calculated as the mean value of the ROI of the lowest ADC value. The same method was applied to a corresponding area in the contralateral normal-appearing white matter judged on both T2WI and contrast-enhanced T1WI. Relative ADC ($rADC$) of the tumors was determined as the ratio of the minimum ADC divided by the mean ADC of the contralateral unaffected white matter. ADC_{min} values were expressed as $\times 10^{-3}$ square millimeters per second.

For assessment of DSC-PWI data, whole-brain CBV maps were generated by applying a single-compartment model and an automated arterial input function. The relative maximum CBV ($rCBV_{max}$) was calculated by dividing the tumor CBV voxel value by the mean CBV value of the contralateral unaffected white matter to minimize variances in $rCBV_{max}$ values in each individual patient. Measurements of $rCBV_{max}$ values were performed with the

same ROIs as those used for ADC measurements. The ROIs for the ADC and $rCBV$ measurements were not identical and were not from the same region of the tumor in each patient. The signal intensities on DWI, $rADC$, ADC_{min} , and $rCBV_{max}$ parameters were determined by another senior neuroradiologist who was experienced in diffusion and perfusion data acquisition and blinded to the tumor histology and molecular data. This method has been shown to provide the highest interobserver and intraobserver reproducibility.²⁹

Immunohistochemistry Staining

Immunohistochemistry was performed on $5\text{-}\mu\text{m}$ -thick sections from paraffin-embedded tumor specimens of all evaluated patients. Sections were incubated overnight at 4°C with the monoclonal anti-IDH1 antibody (DIA-H09; Dianova, Hamburg, Germany) that specifically reacts with the mutant IDH1-R132, the most common glioma-derived mutation,³⁰ but not with the wild-type *IDH1*. Following incubation with horseradish peroxidase–conjugated secondary antibody, the slides were then stained with the Cytomation En-Vision + System horseradish peroxidase (diaminobenzidine) detection

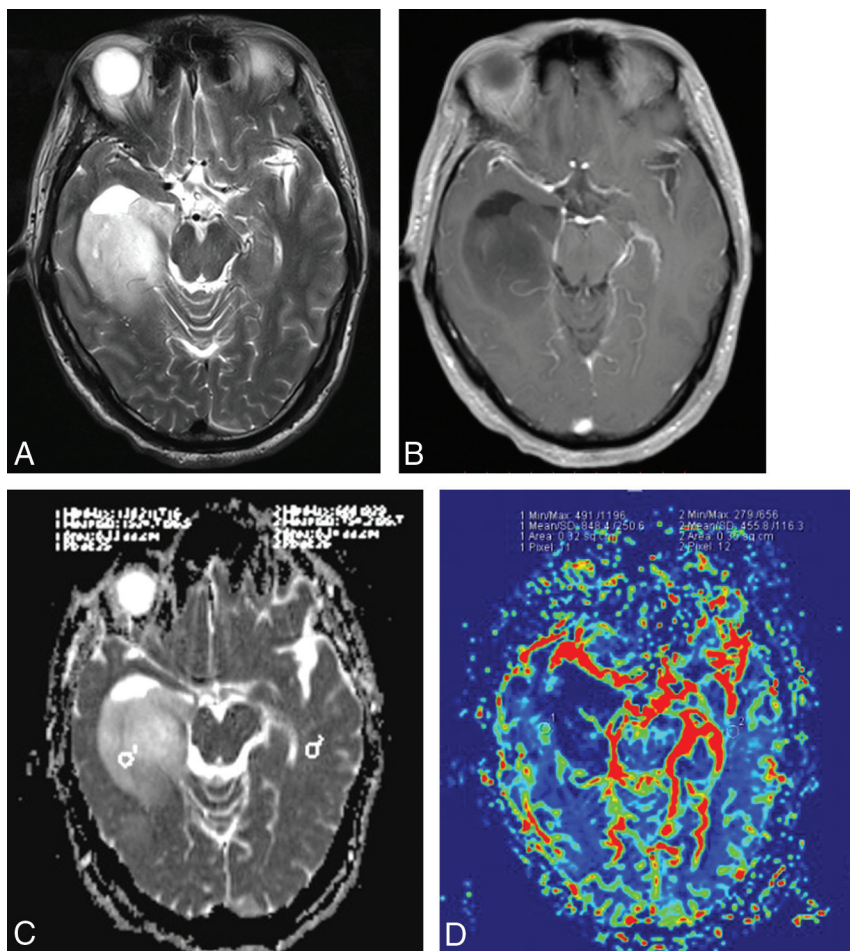


FIG 2. A 50-year-old man with an anaplastic astrocytoma with an *IDH* mutation. A, Axial T2WI demonstrates heterogeneous high signal intensity with sharp borders on the right temporal lobe. B, Contrast-enhanced axial T1-weighted image demonstrates a nonenhancing lesion in the right temporal region. C, A corresponding ADC map shows the tumor with an increased ADC value ($ADC_{min} = 1.456 \times 10^{-3} \text{ mm}^2/\text{s}$, $rADC = 2.51$). D, Correlative color CBV image shows relatively low perfusion with the calculated $rCBV_{max}$ of 1.86.

Table 2: Comparison of DWI and DSC-PWI variables between *IDH* mutation and wild-type grade II and III astrocytomas

	<i>IDH</i> Mutation	<i>IDH</i> Wild Type	<i>P</i> Value
$ADC_{min} (\times 10^{-3} \text{ mm}^2/\text{s})$	1.21 ± 0.27	0.87 ± 0.18	<.001
$rADC$	1.88 ± 0.41	1.37 ± 0.31	<.001
$rCBV_{max}$	1.41 ± 0.50	3.47 ± 2.34	.004

^a Data are means.

kit (Dako, Carpinteria, California) and counterstained with hematoxylin. Staining was interpreted as positive when $\geq 10\%$ of tumor cells showed a strong cytoplasmic staining for mIDH1, whereas staining of $< 10\%$ of tumor cells was counted as negative findings.³¹

DNA Sequencing

Genomic DNA was extracted from formalin-fixed, paraffin-embedded tissue sections by using MygthyAmp for FFPE (Takara Bio, Shiga, Japan), according to the manufacturer's instructions. Mutational alterations of *IDH1* and *IDH2* at hotspot codons R132 and R172 were assessed by a bidirectional cycle sequencing of polymerase chain reaction-amplified fragments with the following primers: *IDH1f* (5'-TGCCACCAACGACCAAGTCA-3') and *IDH1r* (5'-CATGCAAAA TCACATATTTGCC-3'); *IDH2f* (5'-TGAAAGAT-

GGCGGCTGCAGT-3') and *IDH2r* (5'-GGGGTGAAGACCATTGAA-3').

Data Analysis

All quantitative parameters are presented as mean $\pm$ SD. Comparisons of ADC_{min} , $rADC$, and $rCBV_{max}$ values between *IDH*-mutant and wild type of grade II and III astrocytomas were made with the Mann-Whitney *U* test. The receiver operating characteristic and logistic regression analysis were performed to determine the best cutoff value in discriminating *IDH*-mutant from wild-type tumors. The sensitivity, specificity, positive predictive value, negative predictive value, Youden index, and area under the curve based on optimum thresholds for variable parameters were calculated. We chose the cutoff value for each quantitative parameter that provided optimal sensitivity and specificity. In addition, comparisons of the areas under the curve for different variables were made with the *Z*-test. Statistical analysis was calculated in SPSS, Version 19.0 (IBM, Armonk, New York). *P* < .05 was considered significant.

RESULTS

Forty-two histologically confirmed grade II and III astrocytoma cases including 17 cases with *IDH* mutation and 25 cases without such mutation were enrolled in this study. The clinical, histologic, and cMRI characteristics are summarized in Table 1. Twenty-five patients with no *IDH1* or *IDH2* mutations were older (*IDH* mutation = 35.76 ± 9.13 years, *IDH* wild type = 45.96 ± 18.36 years, *P* = .041) and demonstrated more indistinct margins than those with *IDH* mutations (*IDH* mutation, 6/17; *IDH* wild type, 17/25; *P* = .037). Tumors with *IDH* mutations were more likely to occur in the frontal lobes (Figs 1A, -B, and 2A, -B). No differences in heterogeneous appearance were observed among the groups.

The ADC_{min} values, $rADC$ ratios, and $rCBV_{max}$ calculated for *IDH*-mutant and wild-type grade II and III astrocytomas are summarized in Table 2. Both the ADC_{min} (*IDH* mutation = 1.21 ± 0.27 , *IDH* wild type = 0.87 ± 0.18 ; *P* < .001) and $rADC$ (*IDH* mutation = 1.88 ± 0.41 ; *IDH* wild type = 1.37 ± 0.31 ; *P* < .001) were significantly higher in *IDH*-mutant tumors than in wild types (Figs 1C and 2C). The $rCBV_{max}$ (*IDH* mutation = 1.41 ± 0.50 ; *IDH* wild type = 3.47 ± 2.34 ; *P* = .004) in patients with *IDH*-mutant tumors was significantly lower than in those with *IDH* wild-type tumors (Table 2 and Figs 1D and 2D).

The results of the receiver operating characteristic curve analysis are shown in Table 3 and Fig 3. Logistic regression analysis

Table 3: Measurement of TV, sensitivity, specificity, PPV, NPV, YI, and AUC of ADC_{min} values, rADC, rCBV_{max}, cMRI, DWI + DSC-PWI, and cMRI + DWI + DSC-PWI for assessing the IDH status of grade II and III astrocytomas

	TV	Sensitivity (%)	Specificity (%)	PPV (%)	NPV (%)	YI	AUC (95% CI)
ADC _{min}	1.01	76.92	82.61	91.20	60.50	0.60	0.87 (0.71–0.96)
rADC	1.60	84.62	73.91	88.30	67.30	0.59	0.84 (0.68–0.94)
rCBV _{max}	2.35	100.00	60.87	85.60	100.00	0.61	0.82 (0.66–0.93)
cMRI		88.24	52.00	88.50	44.30	0.40	0.78 (0.63–0.89)
DWI + DSC-PWI		100.00	65.22	87.00	100.00	0.65	0.88 (0.54–0.84)
cMRI + DWI + DWI-PWI		92.31	91.30	96.10	83.60	0.84	0.92 (0.78–0.98)

Note:—TV indicates threshold values; PPV, positive predictive value; NPV, negative predictive value; YI, Youden Index; AUC, area under the curve.

was used to test these cMRI parameters among groups; then, a combination of cMRI, DWI, and DSC-PWI for the diagnosis of *IDH* mutation yielded a sensitivity, specificity, positive predictive value, negative predictive value, and Youden index of 92.31%, 91.30%, 96.10%, 83.60%, and 0.84, respectively. A significant difference was found in the areas under the curve between cMRI and cMRI + DWI + DSC-PWI ($Z = 2.8$, $P = .005$).

DISCUSSION

The present study demonstrates that cMRI, DWI, and DSC-PWI can be used to evaluate the *IDH* mutational status in gliomas and that a combination of DWI, DSC-PWI, and cMRI further improves the diagnostic accuracy. We found that patients with *IDH* wild-type tumors were significantly older than those with *IDH*-mutated tumors. This correlation between age and *IDH* mutation status in grade II gliomas has also been reported by Metellus et al.¹⁷ Our finding that *IDH*-mutated tumors tend to reside in the frontal lobes is consistent with previous studies reporting that *IDH*-mutation tumors are strongly associated with frontal locations.^{9,28} In this study, all 11 tumors that did not involve the cerebral cortex were *IDH* wild type, which confirmed a previous finding in anaplastic gliomas that tumors not located in the cerebral cortex were *IDH*-intact tumors.²⁷ While the radiologic appearance of infiltrating lesions in anaplastic gliomas was associated with *IDH* mutation status,¹⁷ these cMRI characteristics lack the ability to quantify the findings. Indeed, it is challenging to determine molecular alterations of such tumors by using cMRI only.

ADC_{min} values have been extensively used to investigate brain tumors and their prognosis.^{21,32–34} Tan et al²⁵ reported that ADC_{min} and rADC could be used to identify gliomas with and without *IDH* mutation. Similarly, Lee et al³⁵ found that the mean ADC value was a useful parameter for differentiating *IDH1* gene mutation–positive high-grade gliomas from the mutation-negative subtypes with histogram analysis. Our study demonstrated that the ADC_{min} and rADC values of *IDH*-mutant grade II and III astrocytomas were higher than those of wild types. ADC_{min} has been shown to depict the sites of highest cellularity within heterogeneous tumors.^{21,33} Therefore, we chose this simple-but-efficient method for the tumor analysis and differentiation. Accumulating evidence has indicated that mutation in the *IDH* gene family could reduce catalytic generation of α -ketoglutarate and, in turn, lead to the production of the oncometabolite (R)-2-hydroxyglutarate [(R)-2HG], ultimately giving rise to increased cell proliferation or cellularity.^{36–38} Therefore, it is conceivable that differences in ADC_{min} and rADC values as presented in the current study might be useful for predicting the molecular profile in astrocytomas with respect to *IDH* mutational status.

DSC-PWI has the potential to noninvasively provide morphologic and functional information about gliomas.^{22,23} Law et al³⁹ reported that DSC-PWI could be used to predict the median time to progression in gliomas and that a lower rCBV corresponds to significantly prolonged progression-free survival. However, these authors did not investigate the rCBV-related molecular mechanisms such as *IDH* mutation. Our data suggest that rCBV_{max} values are significantly associated with the *IDH* mutational status. Recent research showed that *IDH* mutation leads to 2HG (an activator of Egl-9 prolyl-4-hydroxylases) accumulation, resulting in decreased hypoxia-inducible-factor 1- α activation and downstream inhibition of angiogenesis-related signaling.^{38,40} As demonstrated by Kickingeder et al¹⁸ that *IDH*-mutant and wild-type tumors were both associated with distinct imaging phenotypes and were predictable with rCBV imaging in a clinical setting, rCBV maps were used to evaluate *IDH* mutational status of high-grade gliomas with similar results (ie, *IDH* mutant tumors represented considerably lower rCBV).³⁵ Similarly, the rCBV_{max} values in our cases with *IDH* mutation were significantly lower than wild types. Therefore, our findings are in good agreement with prior results and theories. In this study, *IDH*-mutated grade II and III astrocytomas were found to correlate with higher ADC_{min} and rADC and lower rCBV; this correlation corresponds to low levels of cellular density and angiogenesis. These relationships may explain why *IDH* mutation is an independent favorable prognostic marker in patients with gliomas.^{14–19} Glioblastoma is the most common malignant and fatal type of brain tumor, with a poor prognosis.^{5,10} Although the presence of an *IDH* mutation is a strong, independent prognostic factor in gliomas, it had been shown that even *IDH*-mutated glioblastomas exhibited clinical outcomes similar to those of grade III astrocytomas without *IDH* mutation.⁶ The *IDH* mutation is relatively rare in glioblastomas because it is associated with secondary but not primary tumors.⁵ Thus, glioblastomas were excluded from our study.

IDH mutation in diffuse gliomas has been considered the most robust prognostic implication in previous studies.^{10–17} *IDH* mutation was associated with a significantly better clinical outcome with 5-year overall survival (93% compared with 51% for wild type).¹⁷ Although our results demonstrated that MR imaging parameters could noninvasively predict *IDH* mutational status in WHO grade II and III astrocytomas, the correlation between imaging parameters and clinical outcomes has not been studied because of the retrospective nature of this study and a short-term follow-up. Nevertheless, a few studies have demonstrated that low ADC values related directly to poor survival in high-grade astrocytomas and that rCBV values

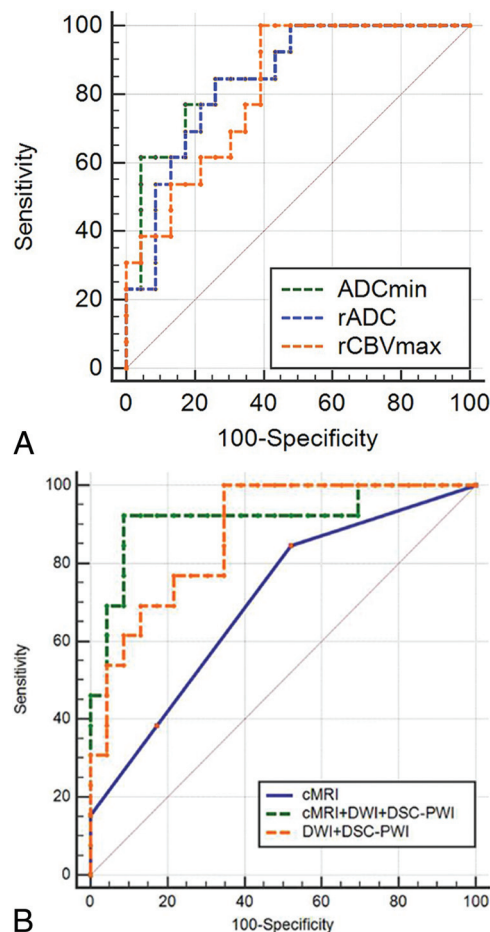


FIG 3. Comparison of receiver operating characteristic curves of ADC_{min} , $rADC$, $rCBV_{max}$ (A) and $cMRI$, $DWI + DSC-PWI$, and $cMRI + DWI + DSC-PWI$ (B) in differentiating *IDH*-mutant grade II and III astrocytomas from wild types.

were positively correlated with median time to progression in patients with gliomas.^{32,39,41,42}

There are a few limitations to this study. It has inherent biases associated with retrospective analyses and a relatively small sample size. A multicentered prospective investigation with a larger sample size is warranted to verify these results and ensure the reproducibility. Second, owing to the short-term follow-up, the clinical outcomes are not available. Imaging parameters as predictors of clinical outcomes should be further studied in larger, prospective trials. Third, the classic biomarker of the human p53 tumor suppressor gene (*TP53*) mutation and O6-methylguanine-DNA methyltransferase (*MGMT*) promoter methylation status was not used in this study for case grouping due to lack of relevant molecular pathologic data in our research. Finally, tumor borders were defined with reference to high signal intensity on T2WI and FLAIR, but it is difficult to differentiate tumor infiltration and peritumoral edema merely on the basis of their radiologic appearance.

CONCLUSIONS

Compared with *IDH* wild-type tumors, *IDH*-mutant tumors tend to have a higher ADC_{min} and $rADC$ and a lower $rCBV_{max}$. Application of $cMRI$ with advanced imaging modalities such as DWI

and $DSC-PWI$ is useful to predict *IDH* mutational status in grade II and III astrocytomas and may provide an important, noninvasive, surrogate marker that should be studied further and clinically correlated in larger, prospective trials.

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